



Versiti Wisconsin, Inc.
638 N 18 St • Milwaukee, WI 53233-2121
414.937.6250 • 800.245.3117 • Fax 414.937.6206
www.versiti.org • labinfo@versiti.org

Patient: **Qual, Mock PNIL**

Birth Date: Gender: Unknown

Patient ID:

Alt ID: POSITIVE

MRN/Req: 3000563537 - 99990000022472

Physician: Not Provided

Additional Copies:

Client #: 9999
ZZ Test Organization
SAMPLE REPORT
638 North 18 Street
Milwaukee, WI 53233-2121

SAMPLE REPORT

Neonatal Alloimmune Thrombocytopenia (NAIT)

Interpretation

Results: No platelet specific alloantibodies detected in maternal serum.

Antibody Specificity: Not applicable.

Strong positive reactions detected in maternal serum against Class I HLA.

Strong positive reactivity detected in maternal serum against intact platelets indicating the presence of platelet-reactive antibodies and/or immune complexes. This reactivity is probably due to HLA antigens. Such antibodies are rarely, if ever, implicated in the pathogenesis of Neonatal Alloimmune Thrombocytopenia (NAIT).

Methodology:

The serologic results were determined using a diverse panel of tests including monoclonal antibody immobilization of glycoproteins and whole platelet flow cytometry assays. The panel is designed to identify platelet antibodies reactive with Class I HLA, as well as antigens on platelet glycoproteins Ia, IIa, IIb, IIIa, Ib/IX, and glycoprotein IV (GPIV/CD36).

This test was developed and its performance characteristics validated by BloodCenter of Wisconsin. It has not been cleared or approved by the FDA. However, the FDA has determined that such clearance or approval is not necessary. The results are not intended to be used as the sole means for clinical diagnosis or patient management decisions but should not be regarded as investigational or for research. Our laboratory is certified under the Clinical Laboratory Improvement Amendments (CLIA) as qualified to perform high complexity clinical laboratory testing.

Date Received: 10/12/2018 1:35 PM

Electronically Signed By:
Ruchika Sharma, MD
PNL Medical Director
Report Date: 10/16/18

Thrombocytopenia Evaluation

Platelet Antibody Screen

Procedure	Result	Collected Date/Time	Received Date	Completed Date
Target 1 -IgG Result	Positive^A	10/12/2018 00:00	10/12/2018	10/16/2018
Target 1 -IgM Result	Negative	10/12/2018 00:00	10/12/2018	10/16/2018

A = Abnormal

C = Critical

L = Low

H = High

CLIA 52D1009037

4608593

Laboratory Director: Matthew W. Anderson, MD, PhD

Printed: 04/30/2019 11:34

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Procedure	Result	Collected Date/Time	Received Date	Completed Date
Target 2 -IgG Result	Positive^A	10/12/2018 00:00	10/12/2018	10/16/2018
Target 2 -IgM Result	Negative	10/12/2018 00:00	10/12/2018	10/16/2018

Platelet Antibody Bead Array (PABA)

Procedure	Result	Collected Date/Time	Received Date	Completed Date
GP IIb/IIIa-HPA1	Negative	10/12/2018 00:00	10/12/2018	10/16/2018
GP IIb/IIIa-HPA3	Negative	10/12/2018 00:00	10/12/2018	10/16/2018
GP Ib/IX-HPA2	Negative	10/12/2018 00:00	10/12/2018	10/16/2018
GP Ia/IIa-HPA5	Negative	10/12/2018 00:00	10/12/2018	10/16/2018
HLA-Class I	Positive	10/12/2018 00:00	10/12/2018	10/16/2018
CD36-GPIV	Negative	10/12/2018 00:00	10/12/2018	10/16/2018

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